# **EXHIBIT E**



Tension-free Support for Incontinence

# GYNECARE TVT™

### Tension-free Vaginal Tape

GYNECARE TVT" Single Use Device
GYNECARE TVT" Reusable Introducer
GYNECARE TVT" Reusable Rigid Catheter Guide
GYNECARE TVT" Reusable Rigid Catheter Guide
GYNECARE TVT" anordning til engangsbrug
GYNECARE TVT" sindfører til flergangsbrug
GYNECARE TVT" stift guiding kateter til flergangsbrug
GYNECARE TVT" hulpmiddel voor eenmalig gebruik
GYNECARE TVT" herbruikbare introducer
GYNECARE TVT" herbruikbare starre kathetervoerder

GYNECARE TVT" -laite, kertakäyttöinen GYNECARE TVT" -sisäänviejä, kestokäyttöinen GYNECARE TVT" -katetrinohjain, kestokäyttöinen, jäykkä Dispositif GYNECARE TVT" à usage unique Introducteur réutilisable GYNECARE TVT" Guide de sonde rigide reutilisable GYNECARE TVT"

GYNECARE TVT\*\* Einmal-Implantat GYNECARE TVT\*\* wiederverwendbares Einführungsinstrument GYNECARE TVT\*\* wiederverwendbare starre Katheterführung

Συσκευή μιας χρήσης GYNECARE TVT" Επαναχρησιμοποιήσιμος εισαγωγέας GYNECARE TVT" Επαναχρησιμοποιήσιμος ακαμπτος οδηγός καθετήρα GYNECARE TVT"

Dispositivo monouso GYNECARE TYT™ Introduttore riutilizzabile GYNECARE TYT™ Guida rigida riutilizzabile per catetere GYNECARE TYT™

Dispositivo de utilização única GYNECARE TVT\*\* Introdutor reutilizável GYNECARE TVT'\* Guia rígido de cateter reutilizável GYNECARE TVT'\*

Sistema para un solo uso GYNECARE TYT™ Introductor reutilizable GYNECARE TVT™ Guía de catéter rígida reutilizable GYNECARE TVT™

GYNECARE TVT™ produkt för engångsbruk GYNECARE TVT™ återanvändbar införare GYNECARE TVT™ återanvändbar stel kateterguide



OEMP15506 LAB0012841v5 01/2015 Made in Switzerland © Ethicon, Inc. 2009

Ethicon, Inc. Route 22 West, P.O. Box 151 Somerville, New Jersey 08876-0151 USA 1-877-ETHICON +1-513-337-6928



GYNECARE TYT™ Single Use Device GYNECARE TYT™ Reusable Introducer GYNECARE TYT™ Reusable Rigid Catheter Guide

Please read all information carefully. Failure to properly follow instructions may result in improper functioning of the device and lead to injury.

Important:
This package insert is designed to provide instructions for use of the Tension-free
Yaginal Tape single use device, Reusable Introducer, and Reusable Rigid Catheter
Guide. It is not a comprehensive reference to surgical technique for correcting Stress
Urinary Incontinence (SUI). The device should be used only by physicians trained in
the surgical treatment of Stress Urinary Incontinence and specifically in implanting
the GYNECARE TVT\*\* Device. These instructions are recommended for general use of the device. Variations in use may occur in specific procedures due to individual technique and patient anatomy.

DESCRIPTION (System)
GYNECARE IVT consists of the following:
GYNECARE IVT's single Use Device, provided sterile (available separately)
GYNECARE IVT'® Reusable Introducer, provided non-sterile (available

separately)
GYNECARETYT'\* Reusable Rigid Catheter Guide, provided non-sterile

(available separately)

GYNECARE TVT DEVICE

The GYIECARE TVT DEVICE

The GYIECARE TVT Device is a sterile single use device, consisting of one piece of undyed or blue (Phthalocyanine blue, color index. Number 74160) PROLERE\*

Polypropylene Mesh (tape) approximately 1/2 x 18 inches (1.1 x 45 cm), covered by a plastic shealth cut and overlapping in the middle, and held between two stainless steel needles bonded to the mesh and sheath with plastic collars. The GYNECARE TVD Device is available in either mechanical cut or laser cut versions for the physician's preference. In determine if the GYNECARE TVD Device implant is mechanical or laser cut, consolt the product code on the device packaging; an (I.) at the end of the number indicates the laser cut mesh.

PROLENE Mesh is constructed of lamitted filaments of extruded polypropylene strands identical in composition to that used in PROLENE\* polypropylene nonabsorbable surgical suture. The mesh is approximately 0.027 inches (0.7 mm) thick. This straterial, when used as a suture, has been reported to be non-reactive and to retain its strength indefinitely in clinical use. PROLENE Mesh is knitted by a process which interlinks each fiber junction.

GYNECARE TVT INTRODUCER

OTNECARE TYI INTRODUCER

The GYNECARE TYI INTRODUCER

The GYNECARE TYI Introducer is provided non-sterile and is reusable. The Introducer is made of stainless steel. It consists of two parts, a handle and an inserted threaded metal shaft. The Introducer is intended to facilitate the passage of the GYNECARE TYI.

Device from the vagina to the abdominal skin. It is connected and fixed to the needle, via the threaded end of the shaft, prior to inserting the needle with the tape.

GYNECARE TYT Rigio CATHETER GUIDE
The CYNECARE TYT Rigio Catheter Guide is a non-sterile reusable instrument intended to facilitate the identification of the urethra and the bladder neck during the surgical procedure. It is inserted into a Foley catheter (recommended size 18 French) positioned in the bladder via the urethra. To facilitate insertion, it can be lubricated with gel.

### INDICATIONS

INDICATIONS
The GYNECABETYT Device is intended to be used as a pubourethral sling for treatment of Stress Urinary Incontinence (SUI), for female urinary incontinence resulting from urethral hypermobility and/or intrinsic sphincter deficiency. The GYNECABETY IT introducer and Rigid Catheter Guide are available separately and are intended to facilitate the placement of the GYNECABETYT Device.

### PATIENT FACTORS

Physicians should use their surgical experience and judgment to determine if PROLENE Mesh is appropriate for certain patients. Patient-specific factors may impair wound healing, which may increase the likelihood of adverse reactions.

- The procedure can be carried out under local anesthesia, but it can also be

- INSTRUCTIONS FOR USE
   The procedure can be carried out under local anesthesia, but it can also be performed using regional or general anesthesia.
   Before the patient is prepped and draped, she should be placed in the lithotomy position taking care to avoid hip flexion greater than 60°.
   Insert an 18 French Folgy catheter and leave it to open drainage.
   At the level of the mid urethra, inject a small amount of local anesthesia submuosally to create a space between the vaginal wall and the perimetrial fascia. The extent of dissection required for placement is minimal. Only a small pararethral incision is required over the mid urethra to position the tip of the Needle. Using forceps, grasp the vaginal wall and to position the urethra. Using a small scaple, make a signitial incision about 1.5 cm long starting approximately 1.0 cm cephalad from the urethral meatus. This incision will be positioned over the mid-urethral zone and will allow for subsequent passage of the limplant.
   With a small pair of blunt scissors, make two small paraurethral lateral dissections (approximately 0.5 to 1.0 cm) to accommodate the tips of the Needle.
   Identify the two Needle exit sites, which should be 2-2.5 cm on each side of the midline, immediately above the pubic symphysis. Either mark these sites or, if desired, place two small 3-4 mm transverse stab incisions at the intended exit site. In order to avoid the inferior epigastric vessels it is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit sites be not more than 2.5 cm from the midline. It is important that the exit site is more than 2.5 cm

- needles.

  10. Gently push the tip of the 18 French Foley catheter toward the posterior
- needles.

  10. Gently push the tip of the 18 French Foley catheter toward the posterior lateral wall of the bladder opposite to the intended Needle passage. For example, by pushing toward the patient's left side the bladder will go from a spherical to a spheroid configuration. This moves the bladder away from the back of the pubic symphysis. Additionally, it moves the bladder neck and the urethra to the left as the Needle is passed on the patient's right side, thereby minimizing the risk of bladder perforation.

  11. Hold the Introducer Handle using your dominant hand. Pass the tip of the Needle that is mounted on the GYNECARETVT Introducer, paramethrally through the urogenital diaphragm at the level of the midurethra. Initial insertion of the device is controlled by using the tip of the lindex finger of the non-dominant hand, which is placed in the varjan aunder the anietior vaginal wall, just lateral to the suburethral incision. The curved part of the Needle should rest in the palm of the non-dominant hand. Pass the Needle through the urogenital diaphragm into the retropuble space. During the initial placement into the paramethral dissected space the Needle tip should be oriented horizontally, i.e. in the frontal plane. During passage through the urogenital diaphragm reststance to the passage of the Needle is significantly reduced once it enters the retropubic space.

- 12. At this point, the non-dominant hand is moved from the vagina to the suprapubic exit site. The Needle tip is guided through the retropubic space staying as dose to the back of the pubic symphysis as possible. This is achieved by lowering the ONIECARE TIM Introducer Handle, thereby pressing the Needle tip against the back of the pubic bone.
  2. During a secare through the attemptic space against the Needle in towards the
- tup against the back of the pubic bone.

  3. During passage through the retropibic space aim the Needle tip towards the pre-marked abdominal exit site.

  14. When the needle tip has reached the abdominal incision unscrew the GYHE/CARE I'VI Introducer from the Needle. Before the Implant is pulled into place, remove the 18 French Foley catheter and perform a cystoscopy (70 decree Jens recommended).
- degree lens recommended).

  15. Once bladder integrity is confirmed, pull the Needle upward to bring the Implant out through the abdominal exit site. Clamp the Implant just below the Needle. Cut the Implant between the connection to the Needle and the
- damprocedure is now repeated on the patient's other side while repeating steps 9—15. NOTE: IN ORDER TO MINIMIZE THE RISK OF BLADDER RIJURY, IT IS IMPORTANT THAT THE BLADDER BE DISPLACED TO THE CONTRALATERAL SIDE USING IT
- IT IS INFOUNDED IT HE DELIVER BY CONTINUED IN STEP 10.

  17. The ends of the implant are then pulled upward to bring the implant (sling) loosely, i.e., without tension, under the indiurethra, Adjust the implant so that leakage is reduced, allowing only a few drops of urinary leakage to occur under sters. For this, use patient feedback, i.e. coughing with a full bladder (approximately 300 mt).

  18. Grasp the Implant Sheaths that surround the Implant with champs, taking care not to grasp the Implant. Then individually remove the Implant Sheaths by gently pulling up on the clamps, away from the abdome, one at a time. Io aword putting tension on the Implant, a blunt instrument (iscsors or forceps) should be placed between the urethra and the Implant during removal of the Implant Sheaths.

  19. NOTE: Premature removal of the sheath may make subsequent adjustments difficult.

- Close the skin incisions with suture or surgical skin adhesive.
   Empty the bladder. Following this procedure, postoperative catheterization is not typically required. The patient should be encouraged to try to empty the bladder 2-3 hours after the operation.

# CONTRAINDICATIONS

As with any suspension surgery, this procedure should not be performed in pregnant patients. Additionally, because the PROLENE Mesh will not stretch significantly, it should not be performed in patients with future growth potential including women with plans for future pregnancy.

- with plans for future pregnancy.

  WARNINGS AND PRECAUTIONS

  Do not use GYNECARE TVT in patients who are on anti-coagulation therapy.

  Do not use GYNECARE TVT in a patient who has a urinary tract infection.

  Users should be familiar with surgical technique for bladder next suspensions and should be adequately trained in the GYNECARE TVT implantation procedure before employing the GYNECARE TVT Device. It is important that the tape be located without tension under mid-urethra.

  Acceptable surgical practice should be followed for the procedure as well as for the management of contaminated or infected wounds.

  The procedure should be performed with care to avoid large vessels, nerves, bladder and bowel. Attention to local anatomy and proper passage of needles will minimize risks.
- bladder and bowel. Attention to local anatomy and proper passage of needles will minimize risks.

  Retropublic bleeding may occur post-operatively. Observe for any symptoms or signs before releasing the patient from the hospital.

  Cytoscopy should be performed to confirm bladder integrity or recognize a bladder perforation.

  The Rigid Catheter Guide should be gently pushed into the Foley catheter so that the catheter guide does not extend into the holes of the Foley catheter. When removing the Rigid Catheter Guide, open the handle completely so that the catheter emails properly in place.

  Do not remove the plastic sheath until the tape has been properly positioned.

- Ensure that the tape is placed with minimal tension under mid-urethra.
   PROLENE Mesh in contaminated areas should be used with the understanding that subsequent infection may require removal of the material.
   The patient should be counseled that future pregnancies may negate
- the effects of the surgical procedure and the patient may again become incontinent.
- incontinent.

  Since no dinical experience is available with vaginal delivery following the procedure, in case of pregnancy delivery via cesarean section is recommended. Post-operatively, the patient is recommended to refrain from heavy lifting and/or exercise (i.e., cycling, jogging) for at least three to four weeks and interocuse for one month. The patient can return to other normal activity
- intercourse for one month. The patient can return to other normal activity after one or two weeks. 
  Should dysura, bleeding or other problems occur, the patient is instructed to contact the surgeon immediately. 
  All surgical instruments are subject to wear and damage under normal use. 
  Before use, the instrument should be visually inspected. Defective instruments or instruments or instruments of instruments of instruments of instruments of instruments of instruments with appear to be corroded should not be used and should be discarded. 
  As with other incontinence procedures, de novo detrusor instability may occur following the procedure. Io minimize this risk, make sure to place the tape tension-free in the mid-urethral position. 
  Do not contact the PROLENE Mesh with any staples, clips or clamps, as mechanical damage to the mesh may occur. 
  Do not resterilize/reuse. Reuse of this device (or portions of this device) may create a risk of product degradation, which may result in device failure

- may create a risk of product degradation, which may result in device failure and/or cross-contamination, which may lead to infection or transmission of blood-borne pathogens to patients and users.

  Discard opened, unused devices.

- Discard opertee, universe and ADVERSE REACTIONS
   Punctures or lacerations of vessels, nerves, structures or organs, including the bladder, urethra or bowel, may occur and may require surgical repair.
   Iransitory local irritation at the wound site may occur.
   As with any implant, a foreign body response may occur. This response could result in extrusion, erosion, exposure, fistula formation and/or inflammation.
- Mesh extrusion, exposure, or erosion into the vagina or other structures or
- Mean extrusion, exposure, or erosion into the vagina or other structures or organs.

  As with all surgical procedures, there is a risk of infection. As with all foreign bodies, PROLENE Mesh may potentiate an existing infection.

  Over correction, i.e., too much tension applied to the tape may cause temporary or permanent lower urinary tract obstruction.

  Acute and/or chronic pain Voiding dysfunction

  Pain with intercourse which in some patients may not resolve.

  Neuronuscular problems, including acute and/or chronic pain in the groin, thigh, leg, pelvic and/or abdominal area may occur.

  Recurrence of incontinence

- Recurrence of incontinence

- Recurrence of incontinence Bleeding including hemorrhage, or hematoma. One or more revision surgeries may be necessary to treat these adverse reactions. PROLENE Mesh is a permanent implant that integrates into the tissue. In cases in which the PROLENE Mesh needs to be removed in part or whole, significant dissection may be required.

### OTHER ADVERSE REACTIONS

- HER ADVENSE REACTIONS
  Seroma
  Urge incontinence
  Urinary frequency
  Urinary retention
  Adhesion formation
  Atypical vaginal discharge
  Exposed mesh may cause pain or discomfort to the patient's partner during
  intercourse.

ACTIONS

Animal studies show that implantation of PROLENE Mesh elicits a minimal inflammatory reaction in tissues and stimulates the deposition of a thin fibrous layer of tissue that can grow through the interstices of the mesh, thus incorporating the mesh mind adjacent tissue. The material is not absorbed, nor is it subject to degradation or weakening by the action of tissue enzymes.

# INSTRUCTIONS FOR CLEANING REUSABLE INSTRUMENTS (GYNECARE TYT

INSTRUCTIONS FOR CLEANING REUSABLE INSTRUMENTS (GYNECARE IVI Introducer, OYNECARE TVT Rigid Catheter Guide)

To ensure the reliability and functionality of the GYNECARE TVI Introducer and 
OYNECARE TVT Region Catheter Guide, clean the instruments before initial use and 
after each procedure. The following are suggested manual and automated methods 
for cleaning the instruments. Prior to cleaning, the GYNECARE TVT Introducer should 
be separated into its component parts (handle and threaded shaft). The Introducer is 
reassembled after cleaning and before sterilization.

### Manual Method:

- Manual Method:

  1. Soak the instrument components in an enzyme cleaner suitable for stainless steel instruments.

  2. Wash in a surgical detergent and disinfecting solution at a temperature of 86°F to 95°F 60°C to 35°C). Remove any contamination from body fluids or tissues using a soft brush.

  3. Place the instrument components in an ultrasonic bath with fresh detergent solution for approximately 10 minutes or follow the instructions below if using an automatic washing cycle.

  4. Rinse thoroughly in a stream of fresh tap water followed by towel drying. The instrument components may be treated with instrument lutricant.

instrument components may be treated with instrument lubricant.

Automated Method:

Automated washing cycles are suitable for stainless steel instruments. One recommended cycle is described below:

Rinse-Rec (yels cold Water — I minute

Wash 176°F (80°C) — 12 minutes

Rinse Cycle — 12 minutes

Rinse Cycle — 12 minutes

Final Rinse — 2 minutes

Final Rinse — 2 minutes

Binse with Pennineralized water 176°F (80°C) — 2 minutes

Dry 199.4°F (93°C) — 10 minutes

Dry 199.4°F (93°C) – 10 minutes
 STERILIZATION RECOMMENDATIONS FOR REUSABLE INSTRUMENTS (GYNECARE TVT Introducer, GYNECARE TVT Rigid Catheter Guide)
 The GYNECARE TVT Introducer and GYNECARE TVT Rigid Catheter Guide are supplied non-sterile. To sertilize, steam autodave prior to each use. Steam anticaive at a temperature of 270°F to 284°F (132°C to 140°C) for a minimum of 4 minutes (pre-vacuum). It is the responsibility of the end user to assure sterility of the product when using sterilization process recommended, since bioburden and sterilization equipment will vary.

## INSTRUMENT MAINTENANCE

INSTRUMENT MAINT LEARNES

- GYNECARE TVT Introducer

Before each use, inspect the threaded parts of the inner shaft.

- GYNECARE TVT Rigid Catheter Guide

Before each use, inspect the instrument. Check to ensure that the long end which traverses the catheter channel has no sharp edges or burrs.

HOW SUPPLIED

The GYNECARE TVT Device is provided sterile (ethylene oxide) for single use. Do not re-sterilize. Do not use if package is opened or damaged. Discard opened, unused devices. The reusable GYNECARE TVT Introducer and GYNECARE TVT Rigid Catheter Guide are supplied separately and are non-sterile. These accessories are to be cleaned and sterilized prior to each use as described above.

### STORAGE

STORAGE

No special storage conditions required. Do not use after expiration date.

Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a licensed healthcare practitioner.

